1	ENGROSSED HOUSE		
	BILL NO. 2676 By: Marti and Davis of the		
2	House		
3	and		
4	Weaver and Bullard of the Senate		
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8	An Act relating to public health and safety; amending 63 O.S. 2011, Section 2-309, as last amended by		
9	Section 1, Chapter 255, O.S.L. 2018 (63 O.S. Supp. 2020, Section 2-309), which relates to the Uniform		
10	Controlled Dangerous Substances Act; updating name of agency; exempting certain practitioners from		
11	electronic prescription requirements for controlled dangerous substances; allowing for the utilization of		
12	electronic prescriptions under certain circumstances; modifying internal statutory references; requiring		
13	practitioners to register with certain agency in order to purchase prescription forms; removing fee		
14	exemption and time period for valid registrations; directing practitioners to purchase prescription		
15	forms from list of approved vendors; allowing certain content to be included on prescription forms;		
16 17	expanding definition to allow for the inclusion of certain information on electronic prescriptions;		
17	clarifying authority of licensed practitioners to purchase prescription forms; and providing an		
10	effective date.		
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20	BE IT ENACTED BY THE PEOPLE OF THE STATE OF OKLAHOMA:		
22	SECTION 1. AMENDATORY 63 O.S. 2011, Section 2-309, as		
23	last amended by Section 1, Chapter 255, O.S.L. 2018 (63 O.S. Supp.		
24	2020, Section 2-309), is amended to read as follows:		

1 Section 2-309. A. 1. Except for dosages medically required 2 for a period not to exceed forty-eight (48) hours which are 3 administered by or on direction of a practitioner, other than a 4 pharmacist, or medication dispensed directly by a practitioner, 5 other than a pharmacist, to an ultimate user, no controlled dangerous substance included in Schedule II, which is a prescription 6 7 drug as determined under regulation promulgated by the State Board of Pharmacy, shall be dispensed without an electronic prescription 8 9 of a practitioner; provided, that in emergency situations, as 10 prescribed by the State Board of Pharmacy by regulation, such drug 11 may be dispensed upon oral prescription reduced promptly to writing 12 and filed by the pharmacist in a manner to be prescribed by rules 13 and regulations of the Director of the Oklahoma State Bureau of 14 Narcotics and Dangerous Drugs Control.

15 2. Electronic prescribing shall be utilized for Schedules II,
16 III, IV, and V, subject to the requirements set forth in 21 CFR,
17 Section 1311 et seq.

3. An electronic prescription with electronic signature may
serve as an original prescription, subject to the requirements set
forth in 21 CFR, Section 1311 et seq.

4. Prescriptions shall be retained in conformity with the
requirements of this section and Section 2-307 of this title. No
prescription for a Schedule II substance may be refilled.

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1 5. The electronic prescription requirement provided for in this 2 section shall not apply to prescriptions for controlled dangerous 3 substances issued by any of the following: 4 a person licensed to practice veterinary medicine, a. 5 b. a practitioner who experiences temporary technological or electrical failure or other extenuating 6 7 circumstance that prevents the prescription from being transmitted electronically; provided, however, that 8 9 the practitioner documents the reason for this 10 exception in the medical record of the patient, 11 с. a practitioner, other than a pharmacist, who dispenses 12 directly to an ultimate user, 13 d. a practitioner who orders a controlled dangerous 14 substance to be administered through an on-site 15 pharmacy in: 16 a hospital as defined in Section 1-701 of this (1)17 title, 18 (2) a nursing facility as defined in Section 1-1902 19 of this title, 20 a hospice inpatient facility as defined in (3) 21 Section 1-860.2 of this title, 22 (4) an outpatient dialysis facility, 23 a continuum of care facility as defined in (5) 24 Section 1-890.2 of this title, or

1		(6) a penal institution listed in Section 509 of
2		Title 57 of the Oklahoma Statutes,
3	e.	a practitioner who writes a prescription to be
4		dispensed by a pharmacy located on federal property,
5		provided the practitioner documents the reason for
6		this exception in the medical record of the patient,
7		or
8	f.	a practitioner that has received a waiver or extension
9		from his or her licensing board <u>,</u>
10	<u>g.</u>	a practitioner who prescribes a controlled dangerous
11		substance for a supply that when taken as prescribed
12		would be consumed within seventy-two (72) hours, or
13	h.	a practitioner who determines that an electronic
14		prescription cannot be issued in a timely manner and
15		the condition of the patient is at risk.
16	6. Elect	ronic prescriptions shall not $\underline{may}$ be utilized under the
17	following cir	cumstances:
18	a.	compound compounded prescriptions containing two or
19		more commercially available products or two or more
20		active pharmaceutical ingredients,
21	b.	compounded infusion prescriptions containing two or
22		more commercially available products or two or more
23		active pharmaceutical ingredients, or
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- c. prescriptions issued under approved research protocols<del>, or</del>
- d. if the practitioner determines that an electronic prescription cannot be issued in a timely manner and the condition of the patient is at risk.

7. A pharmacist who receives a written, oral or facsimile
prescription shall not be required to verify that the prescription
falls under one of the exceptions provided for in paragraph 6 of
this subsection. Pharmacists may continue to dispense medications
from otherwise valid written, oral or facsimile prescriptions that
are consistent with the provisions of this act.

8. Practitioners shall indicate in the health record of a patient that an exception to the electronic prescription requirement was utilized.

9. All prescriptions issued pursuant to paragraphs paragraph 5
and <u>subparagraph c of paragraph</u> 6 of this subsection shall be <del>issued</del>
on an official prescription form <del>provided</del> <u>approved</u> by the Oklahoma
State Bureau of Narcotics and Dangerous Drugs Control.

1910. a.Effective January 1, 2020, practitioners<br/>PractitionersPractitioners<br/>Practitioners20shall register be registered<br/>Bureau of Narcotics and Dangerous Drugs Control in<br/>order to be issued purchase<br/>official prescription21order to be issued purchase<br/>forms. Such registration shall include, but not be<br/>limited to, the primary address and the address of

each place of business to be imprinted on official prescription forms. Any change to a registered practitioner's registered address shall be promptly reported to the practitioner's licensing board and the Bureau by the practitioner in a manner approved by the Bureau.

- 7 A practitioner's registration shall be without fee and b. subject to approval by the Bureau. Such registration 8 9 shall be valid for a period of two (2) years and may 10 be denied, suspended or revoked by the Bureau upon a 11 finding by the Bureau or licensing board that the 12 registered practitioner has had any license to 13 practice a medical profession revoked or suspended by 14 any state or federal agency.
- 15 Where the Bureau has revoked the registration of a <del>c.</del> 16 registered practitioner, the Bureau may revoke or 17 cancel any official prescription forms in the 18 possession of the registered practitioner. Any 19 revocation or any suspension shall require the 20 registered practitioner to return all unused official 21 prescription forms to the Bureau within fifteen (15) 22 calendar days after the date of the written 23 notification.

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1 A practitioner that has had any license to practice <del>d.</del> c. 2 terminated, revoked or suspended by a state or federal 3 agency may, upon restoration of such license or 4 certificate, register to be issued official 5 prescription forms with the Bureau. Except as provided in subparagraph f of this 6 11. a. 7 paragraph, the Bureau shall issue official Official prescription forms free of charge only to registered 8 9 practitioners in this state. Such forms shall not be 10 transferable. The number of official prescription 11 forms issued to a registered shall be purchased at the 12 expense of the practitioner at any time shall be at 13 the discretion of or the employer of the practitioner 14 from a list of vendors approved by the Bureau. 15 Official prescription forms issued to a registered b. 16 practitioner shall be imprinted only with the primary 17 address and may include other addresses listed on the 18 registration of the practitioner to identify the place 19 of origin. Such prescriptions shall be sent only to 20 the primary address of the registered practitioner. 21 Official prescription forms issued to of a registered с. 22 practitioner shall be used only by the practitioner to 23 whom they are issued designated on the official 24 prescription form.

- d. The Bureau may revoke or cancel official prescription
  forms in possession of registered practitioners when
  the license of such practitioner is suspended,
  terminated or revoked.
- e. Official prescription forms of registered
  practitioners who are deceased or who no longer
  prescribe shall be returned to the Bureau at a
  designated address. If the registered practitioner is
  deceased, it is the responsibility of the registered
  practitioner's estate or lawful designee to return
  such forms.
- 12 f. The Bureau may issue official prescription forms to 13 employees or agents of the Bureau and other government 14 agencies for the purpose of preventing, identifying, 15 investigating and prosecuting unacceptable or illegal 16 practices by providers and other persons and assisting 17 in the recovery of overpayments under any program 18 operated by the state or paid for with state funds. 19 Such prescription forms shall be issued for this 20 purpose only to individuals who are authorized to 21 conduct investigations on behalf of the Bureau or 22 other government agencies as part of their official 23 duties. Individuals and agencies receiving such 24 prescription forms for this purpose shall provide

1 appropriate assurances to the Bureau that adequate 2 safeguards and security measures are in place to 3 prevent the use of such prescription forms for 4 anything other than official government purposes. 5 12. a. Adequate safeguards and security measures shall be undertaken by registered practitioners holding 6 7 official prescription forms to assure against the loss, destruction, theft or unauthorized use of the 8 9 forms. Registered practitioners shall maintain a 10 sufficient but not excessive supply of such forms in 11 reserve.

- 12 b. Registered practitioners shall immediately notify the 13 Bureau, in a manner designated by the Bureau, upon 14 their knowledge of the loss, destruction, theft or 15 unauthorized use of any official prescription forms 16 issued to them, as well as the failure to receive 17 official prescription forms within a reasonable time 18 after ordering them from the vendor approved by the 19 Bureau.
- c. Registered practitioners shall immediately notify the
   Bureau upon their knowledge of any diversion or
   suspected diversion of drugs pursuant to the loss,
   theft or unauthorized use of prescriptions.

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1 B. 1. Except for dosages medically required for a period not 2 to exceed seventy-two (72) hours which are administered by or on direction of a practitioner, other than a pharmacist, or medication 3 4 dispensed directly by a practitioner, other than a pharmacist, to an 5 ultimate user, or the circumstances provided for in paragraphs 5 and 6 of subsection A of this section, no controlled dangerous substance 6 7 included in Schedule III or IV, which is a prescription drug as determined under regulation promulgated by the State Board of 8 9 Pharmacy, shall be dispensed without an electronic prescription. 10 2. Any prescription for a controlled dangerous substance in

11 Schedule III, IV or V may not be filled or refilled more than six 12 (6) months after the date thereof or be refilled more than five 13 times after the date of the prescription, unless renewed by the 14 practitioner.

C. Whenever it appears to the Director of the Oklahoma State Bureau of Narcotics and Dangerous Drugs Control that a drug not considered to be a prescription drug under existing state law or regulation of the <u>State</u> Board of Pharmacy should be so considered because of its abuse potential, the Director shall so advise the <u>State</u> Board of Pharmacy and furnish to the Board all available data relevant thereto.

D. 1. "Prescription", as used in this section, means a written, oral or electronic order by a practitioner to a pharmacist for a controlled dangerous substance for a particular patient, which

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1 specifies the date of its issue, and the full name and address of 2 the patient and, if the controlled dangerous substance is prescribed 3 for an animal, the species of the animal, the name and quantity of 4 the controlled dangerous substance prescribed, the directions for 5 use, the name and address of the owner of the animal and, if written, the signature of the practitioner. When electronically 6 prescribed, the full name of the patient may include the name and 7 8 species of the animal.

9 2. "Registered practitioner", as used in this section, means a 10 licensed practitioner duly registered with the Oklahoma State Bureau 11 of Narcotics and Dangerous Drugs Control <u>authorized</u> to <del>be issued</del> 12 purchase official prescription forms.

E. No person shall solicit, dispense, receive or deliver any controlled dangerous substance through the mail, unless the ultimate user is personally known to the practitioner and circumstances clearly indicate such method of delivery is in the best interest of the health and welfare of the ultimate user.

18 SECTION 2. This act shall become effective November 1, 2021.
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1	Passed the House of Representatives the 8th day of March, 2021.
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4	Presiding Officer of the House of Representatives
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6	Passed the Senate the day of, 2021.
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8	Presiding Officer of the Senate
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